

INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:
H01Q 13/24, A61N 5/04

(11) International Publication Number: WO 95/04385

(43) International Publication Date: 9 February 1995 (09.02.95)

(21) International Application Number: PCT/GB94/01565

(22) International Filing Date: 19 July 1994 (19.07.94)

(30) Priority Data:
9315473.0
9401912.2
27 July 1993 (27.07.93)
1 February 1994 (01.02.94)
GB

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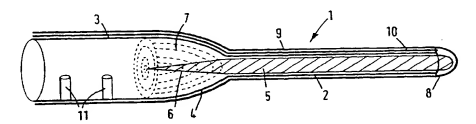
(81) Designated States: AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, ES, FI, GB, GE, HU, JP, KE, KG, KP, KR, KZ, LK, LT, LU, LV, MD, MG, MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SI, SK, TJ, TT, UA, US, UZ, VN, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG), ARIPO patent (KE, MW, SD).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: TREATMENT APPARATUS



(57) Abstract

A probe (1) is designed to propagate and radiate microwave electromagnetic energy in a controlled fashion. The probe (1) includes at least one waveguide (2) of cross-section which would not normally pass microwaves at the operational frequency. The waveguide (2) therefore includes dielectric material (5), such as alumina, in the form of a rod an exposed portion of which forms an antenna. The probe is preferably for use in endometrial ablation and therefore the reduced dimension of the waveguide can be made compatible with the narrow neck of the uterus.

is about 5 millimetres thick and covers the whole of the inner wall of the uterus. Menorrhagia can be cured, or at least alleviated, if the endometrium is wholly or partially destroyed without surgery. This destruction can either be achieved by physical means or by heating the tissue or a combination of both. In common with most body tissue, a temperature of around 60°C maintained in the endometrium for up to 5 minutes will destroy its cells. Because it will no longer be possible for the endometrium to regenerate the lining the condition will be cured.

The current known alternative techniques to hysterectomies work with varying degrees of success but all have disadvantages. The uterus is a very delicate V-shaped pouch-like structure and the opposite walls are normally separated by a thin film of fluid or may be partly in contact. Therefore it is difficult to gain access to the endometrium for the purpose of direct physical treatment or for heating it. It is particularly difficult to treat the tissue immediately surrounding the entrance as heating must be confined to the endometrium itself and not extend to the main body of the uterus and beyond it.

The easiest and least complicated alternative method uses a steel ball about 5mm in diameter heated

by a monopolar connection to a power supply. The ball is rolled around in the uterus under the control of the surgeon to destroy the endometrium. However, the method is time consuming and requires highly specialised surgical experience. Even in skilled hands localised burning can occur or other areas are not fully treated.

It is also known to use certain forms of electromagnetic energy, for example, cell destruction has been achieved by laser ablation where light waves are used. However, laser treatment requires expensive laser equipment and the treatment has to be carried out using highly specialised surgical skills.

From European Patent Publication 0407057 it is known to use radio frequency electromagnetic energy. For example, the method disclosed in that patent involves placing a radio frequency probe in the uterus and setting up a radio frequency field between it and a steel belt around the patients waist. The treatment takes up to 45 minutes including anaesthetic induction and recovery. The procedure itself takes about 15-20 minutes and requires the full time attention of a skilled gynaecologist in moving the probe. This is because, as the typical power used is about 550 watts and radio frequency electromagnetic radiation is difficult to contain it

has to be moved close to the endometrium to be at all effective. It also has the disadvantage that radio frequency electromagnetic energy readily passes through most materials (including tissue) and may very easily leak and insidiously cause injury to both the patient and surgical staff during the course of treatment.

Another method using radio frequency energy is disclosed in European Patent Publication No. 0115420 which discloses a device for hyperthermia therapy using first and second electrodes at a frequency of about 3-30 MHZ.

An object of the present invention is to provide an improved apparatus and method using microwave frequency electromagnetic radiation.

Microwaves at about 2.7GHz are commonly used for cooking because of the strong absorption of radiation at that frequency by water. It therefore might be thought that given the use of light frequency and radio frequency electromagnetic radiation it would be obvious to try microwaves. However, there are no particular restrictions on waveguide or cavity size with microwave ovens therefore a frequency as low as 2.7GHz and a wavelength of 100cm or more presents no problem.

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However, the neck of the uterus can only be dilated to about 10mm diameter at maximum and any probe therefore needs to have a diameter of no more than 8mm for general use. With conventional design this would mean that the microwave frequency would need to be much too high using conventional waveguide of these dimensions and not enough power would be delivered to the endometrium i.e. the higher the frequency the less the depth of absorption by the tissue being treated. It was also thought that standing wave patterns would produce non-uniform heating. However, standing waves only occur where there are reflections present and we have found that the walls of the uterus rather than causing any reflections absorb the waves by the lossy tissue of the endometrium which rapidly reduces the wave to zero before it can reach any potential reflecting objects.

According to the present invention there is provided a probe for applying electromagnetic radiation at microwave frequency to a body comprising means for receiving microwave signal input of a predetermined frequency, a waveguide for receiving and propagating said microwave frequency input, said waveguide being of a cross-sectional dimension which would not normally pass the microwaves at said

frequency, dielectric material within the waveguide, the dielectric constant of which varies the cut-off frequency of the waveguide so that it may propagate desired modes of the microwaves at said predetermined frequency, and an exposed antenna portion at or adjacent an end of the probe allowing wave transmission away from the probe.

The means for receiving the microwave signal may comprise a second waveguide, transition means being provided between the first and second waveguides. In this arrangement the first waveguide is suitably a circular waveguide typically of about 10mm diameter. The second waveguide may also be a circular waveguide of about 20mm diameter. The transition means comprises a tapered waveguide interconnecting the first and second waveguides and loaded with dielectric material.

The dielectric material is preferably in the form of a ceramic rod having a tapered end at the transition to optimise transition and extending outwardly beyond the first waveguide to form the exposed antenna portion of the probe. The use of a dielectric filled first waveguide in accordance with the invention allows the first waveguide to be of smaller diameter because, at a given frequency, the wavelength in dielectric is shorter. Hence, the

diameter of the probe in wavelengths remains constant throughout transition. For any given wavelength the minimum diameter of the probe is around one half of a wavelength. Any smaller and the wave will not pass through. The tapered end of the ceramic rod overcomes the dielectric mismatch between air in the second waveguide and the ceramic material. Without the taper there would be a danger of a reflection at the interface between the first and second waveguides.

In an alternative arrangement a single waveguide is provided and the means for receiving the microwave input directly excites the dielectric filled waveguide of the desired smaller cross-sectional dimension.

The preferred form of probe includes temperature sensors disposed between the first waveguide and a protective sheath. The sensors may be of different lengths in order to detect temperatures at different locations along the length of the probe and are united at a temperature sensor interface.

Although it is preferred that the probe be a single unit it is possible for the probe to comprise two or more separable portions. Therefore, according to another aspect of the invention a probe for applying electromagnetic radiation at microwave

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frequency to a body has a first dielectric stage and a second dielectric stage, the two stages, in use, being operatively connected together the first dielectric stage comprising a first waveguide of a first cross-section; a second waveguide of a second cross-section larger than the cross-section of the first waveguide for receiving and propagating microwave signal input of a predetermined frequency, and transition means between the first and second waveguides including dielectric material, the dielectric constant of which varies the cut-off frequency of the first waveguide so that it may propagate said microwave signal at the predetermined frequency; and, the second dielectric stage comprising a probe antenna of dielectric material, a third waveguide about a portion of the dielectric material and being of substantially the same crosssection as the first waveguide, and an exposed antenna portion at or adjacent a free end of the probe allowing wave transmission away from the probe.

Preferably, the transition means of the first dielectric stage comprises a tapered waveguide interconnecting the first and second waveguides, a tapered end on the dielectric material within the tapered waveguide to optimize transition and a dielectric buffer between the tapered end of

dielectric material and the tapered waveguide, the dielectric constant of which is greater than air but less than that of the dielectric material.

In this arrangement the probe may be for endometrial ablation and the second dielectric stage may include opposed inflatable catheters to aid positioning in the uterus. Suitably, the second dielectric stage also includes temperature sensing means. Where provided with two stages the probe includes interface means for the temperature sensing means and for the inflation of the catheters at the connection between the first and second stages of the probe.

If desired, the exposed antenna portion may include guidance means for selective transmission of the microwaves. The guidance means may comprise a thin metallic layer tapering toward the outer end of the exposed antenna portion to equalise leakage of the microwave energy along the length of the exposed portion. The metal may be Chromium which varies in thickness along the length of the rod instigating a differential relationship of wave reflection and transmission, thus radiating power evenly across the cylindrical area of the probe. Alternatively the guidance means could be mesh varying in grading along

the exposed length of the rod or spaced sold rings the spacing between which is gradually increased.

Where the probe is to be used for medical treatment such as endometrial ablation it is important that the probe be sterile for each use. Although it would be possible to provide a disposable probe this is regarded as unnecessarily expensive. Accordingly, preferably the probe includes a removable and disposable sheath which encapsulates the probe during use.

Therefore, according to another aspect of the invention there is provided a protection means for a probe for applying electromagnetic radiation at microwave frequency to a body, said protection means comprising a sheath having a tubular body which may pass over the probe to encapsulate the operative end of the probe and which is substantiality transparent to microwaves at the intended frequency of operation, and means for securing the sheath in position whereby the sheath may be removed and discarded after use of the probe. Preferably the sheath is transparent and the waveguide includes a graticule or measurement marking to aid insertion.

The protection means preferably further includes a disposable handle arranged to receive a probe in use, the handle being locked in position about the

probe by interengagement with the sheath. The protection means suitably includes a unique marking, such as a bar code, to ensure single use. The protective sheath may also include a bar code.

Although the probe and apparatus of the present invention may be used for any desired application it is preferred that the probe be used for endometrial ablation. Therefore, according to the preferred method of the invention there is provided a method of endometrial ablation comprising the steps of providing a probe as aforesaid having at least an operative end of outside dimensions no greater than the dimensions of a dilated cervix, inserting the operative end of the probe through the cervix into the uterus, applying microwave energy to the probe at a frequency which will be substantially completely absorbed by the endometrium, monitoring the operating temperature to ensure that the endometrium tissue is heated to about 60°C and maintaining the application of the microwave energy for a period of time sufficient to destroy the cells of the endometrium. The microwave energy may be applied continually or in pulses.

The use of microwave power to heat the endometrium has two main advantages. Firstly, electromagnetic radiation at microwave frequencies is

strongly absorbed by tissue and at around 8-12GHz all microwave power is absorbed in a layer of tissue about 5mm thick and it is impossible for microwave heating to extend beyond this region. This is ideal for the treatment of the endometrium which is about 5mm thick. Secondly, because of this strong absorption, the amount of power required to achieve the desired temperature is relatively small compared with RF frequencies and it is likely that the necessary energy could be delivered over a much shorter period than other current treatments take. If desired the radiation might be pulsed so that the tissue is momentarily heated above 60°C and the total treatment time could then be shorter still.

The depth of material over which the microwave power is absorbed depends upon frequency and the material electrical properties. To set this to be around 5mm in the endometrial tissue requires a frequency of about 8-12GHz. This frequency then determines the dimensions of the waveguide needed to carry the wave. If a conventional waveguide were used a diameter of around 20mm would be required. This is clearly far too large to enter the uterus. In accordance with the invention cut-off wavelength is effectively reduced by the use of high dielectric constant material such as ceramic material or

plastics dielectric material or other suitable material which provides a transition to a waveguide of outside diameter of about 8mm.

With the probe of the present invention there is no possibility of radiation leakage and inadvertent heating occurring outside of the uterus along the line delivering power to the implanted antenna. The problem of delivering power through the narrow neck has therefore been solved.

Having delivered the power into the uterus, the power is then distributed uniformly into the roughly flat triangular shaped pouch formed by the uterus by means of the exposed portion of the antenna which is arranged to prevent radiation escaping close to the input end. The temperature increase necessary to destroy the cells of the endometrium may require only 60 watts of microwave power to provide a treatment time of 2.5 minutes.

It may be found that access to the inner wall of the uterus is difficult and in such a case, there is an attribute of microwaves which can be used to advantage to provide an even distribution of the heating effect. In particular, microwaves will only be strongly absorbed by tissue and not by any intervening gas. If desired the uterus may be inflated by a gas such as carbon dioxide so that the

walls will be held away from the antenna and receive an even radiation dose. The gas may be supplied through a central bore formed in the ceramic rod/ If the probe includes inflatable catheters then these may be selectively inflated as required to aid insertion and positioning within the uterus. The probe may also be provided with fibre-optic vision if desired.

The invention also includes a system for selective microwave transmission comprising a probe as aforesaid and a source of microwave energy.

Preferably, the variable parameters of the system are computer controlled.

The invention will now be described by way of example with reference to the accompanying drawings in which:

Figure 1 is a diagrammatic side elevation of a preferred probe in accordance with the invention;

Figure 2 is a block diagram of the preferred system incorporating the probe of Figure 1;

Figure 3 is a diagrammatic side elevation of a second embodiment of probe in accordance with the invention;

Figure 4 is a block diagram of the system incorporating the probe of figure 3;

Figure 5 is a diagrammatic side elevation of a third embodiment of probe in accordance with the invention;

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Figure 6 is a diagrammatic side elevation of a fourth embodiment of probe in accordance with the invention;

Figure 7 is a diagrammatic side elevation of a probe in accordance with the invention including a protective sheath;

Figures 8a, 8b and 8c are diagrammatic views of an arrangement for ensuring single use of the protective sheath; and

Figures 9a and 9b are simplified views showing a probe of the present invention in use.

In figure 1 a microwave probe (1) has a first circular waveguide (2) of a first diameter at one end being of custom-determined diameter according to probe use and a second circular waveguide (3) of a second, larger diameter at the other end. The transition between the first waveguide (2) and the larger diameter second waveguide (3) comprises a frusto-conical waveguide (4) and a dielectric rod (5) located mainly within the first waveguide (2). The dielectric rod (5) has a tapered end (6) extending into the transition waveguide (4). Disposed about the dielectric tapered end (6) is a dielectric buffer

plug (7) having dielectric properties greater than air but less than that of the dielectric rod (6).

The first waveguide (2) extends towards the free end of the probe (1) but terminates short of the free end to leave an exposed antenna portion (8). The exposed antenna portion (8) and the first waveguide (2) are provided with a protective removable and disposed sheath (9) of bio-medically inert and microwave transparent material, for example a protective PTFE or similar material, which may be profiled as shown according to probe use. In order to sense the operating temperature, the probe (1) incudes thermocouple wire temperature sensing means (10).

As can be seen from figure 1 the second waveguide (3) also includes waveguide tuning stubs (11). The stubs (11) are set in the wall of the second waveguide (3) to provide means of intrinsically matching the antenna portion (8) in a body. A probe matched to a specific load, preferably endometrium tissue in this application will relieve the need for extensive pre-operative tuning. In addition, the provision of stubs (11) limit the existence of standing waves in the coaxial feed line (12) which can form there when matching is initiated at the system tuning network end of the coaxial feed line.

Standing waves in the coaxial feed line will generate heat and reduce the working life of the cable.

However, subtle load variations from patient to patient can be fine tuned using the system tuning network (13) shown in figure 2. In figure 2, the probe (1) of the invention is supplied with a microwave frequency input in the microwave spectrum, preferably in the region of 8-12GHz, from a microwave frequency generator source and amplifier (14). The amplified signal is passed to the probe (1) via waveguide line (15) and the coaxial feed line (12). Although, the provision of stubs (11) permits the tuning of the probe to the specific load, fine tuning is provided by the tuning network (16) controls the fine turning of the match of power into the loaded probe. The power level of the source/amplification unit (14) is monitored by a power sense (17) on the waveguide line (15). A thermometry unit (18) is provided to take temperature sensor readings at the probe/tissue interface (1). The various signals are collated and conditioned and fed to a PC/user interface (19) which may interface with a user's conventional PC graphics monitor (20). In this way the user may vary the frequency of the source (14), set the power level required, and vary the tuning network (16) to achieve optimum match into a load. Also during the treatment, real-time graphs of temperature data can be viewed on the monitor (20).

In the embodiment of figures 3 and 4 the probe arrangement is similar to that described with reference to figures 1 and 2 except that the probe is formed in two parts. In figure 3 a microwave probe (101) has a dielectric input stage (102) and a dielectric output stage (103). The input stage (102) includes a circular waveguide (104) of a first diameter at one end and a circular waveguide (105) of a second, smaller diameter at the other end, the diameter being of custom-determined diameter according to probe use. The transition between the waveguide (104) and the smaller diameter waveguide (105) comprises a frusto-conical waveguide (106) and a first dielectric rod (107) located mainly within the waveguide (105) but having a tapered end (108) extending into the transition waveguide (106). Disposed about the dielectric tapered end (108) is a dielectric buffer plug (109) having dielectric properties greater than air but less than that of the dielectric rod (107). The circular waveguide (105) terminates in a flange (110) and the rod (107) extends beyond the flange (110) to a joint (111).

The dielectric output stage (103) includes a second dielectric rod (112), an inner end of which abuts the end of the dielectric rod (107 at the joint (111). The output stage (103) is provided with a further waveguide (113) which extends from the flange (110) towards the free end of the probe (101). However, the waveguide (113) terminates short of the free end of the probe (101) to leave an exposed antenna portion (114). The exposed antenna portion (114) and the waveguide (113) are provided with a protective sheath (115) of PTFE or other suitable material as with the first embodiment. In order to sense the operating temperature, the probe (101) includes thermocouple wire temperature sensing means (130). The temperature sensing means (130) is connected to a temperature sensor interface (116) at the flange (110).

The probe (101) disclosed by way of example is a probe for endometrial ablation and, in order to facilitate insertion of the probe inside the uterus, the probe (101) includes two balloon catheters (117) (only one shown), one fixed to each side of the waveguide (113). The catheters (117) are provided with air by means of air tubes (118) and an air tube interface (119) is provided adjacent the flange (110) on the circular waveguide (105).

The probe system of figure 3 is preferably arranged as disclosed in figure 4. In that arrangement, it will be seen that the probe (101) is supplied with a microwave frequency input in the region of 8-12GHz from a microwave frequency generator source (120), the signal of which is amplified by amplifier (121) and passed through a tuning network (122) before entering the input dielectric stage (112) at the circular waveguide (114). The tuning network (122) controls the match of power into a loaded probe (101) and the match is monitored using a power meter (123). Personal computer instrumentation (124) is used to vary the frequency of the source (120), set the power level required, and vary the tuning network (122) to achieve optimum match into a load. This could also be done manually, if required. A thermometry unit (125) is provided to take temperature sensor readings from the probe (101) received via the interface (116) and store these on disk in the p.c. (124). the treatment, real-time graphs of temperature data can be viewed on the monitor (126).

In order to facilitate manipulation of the probe within the uterus, an inflation unit 127 is provided which is operative to supply sufficient air pressure to inflate the catheters (117) on the probe surface.

The probe 140 of the embodiment of figure 5 is similar to that of figure 3 and where appropriate similar references have been used. The main difference in the embodiment of figure is that the waveguide surrounding the dielectric rod (107) is formed by thermocouple wire 142 coiled about the exposed antenna portion 114 for temperature sensing. The flange 110 is again separable into two parts 144,146 each of which includes thermocouple connectors allowing connection of the thermocouple wire 142 to a thermocouple interface 148. In order to serve as a waveguide as well the thermocouple wire 142 is wound so as to provide controlled radiation along the length of the dielectric rod 107.

The embodiment of figure 6 is an alternative arrangement where there is a single waveguide. In this arrangement a microwave probe 201 has a circular waveguide 202 filled with a dielectric material 203. The waveguide 202 terminates short of the end of the probe 201 providing an exposed antenna portion 204. Towards the end of the probe 201 remote from the exposed antenna portion 204 there is a coaxial feed line input 205 and a waveguide excitation stub 206 which directly excites the dielectric filled waveguide 202. The probe 201 is matched to the load of the body into which it is to be inserted by means

of tuning stubs 207 fixed to the wall of the waveguide 202.

As with previous embodiments the probe 201 is provided with a protective sheath 208 of PTFE or other suitable material and reference is particularly directly to the disclosure of one form of the sheath given in figure 7. A temperature sensor 209 is provided between the sheath 208 and the waveguide 202 feeding a temperature indicative signal back to the control (not shown).

In figure 7 an embodiment similar to the embodiments of figures 3-5 is illustrated where the probe 301 includes a first waveguide 302 of small diameter, a second waveguide 303 of larger diameter and a frusto-conical transition waveguide 304 between the two. The first waveguide includes a dielectric rod 305 one end 306 of which is tapered at the transition and the other end of which provides an exposed antenna portion 307. The respective waveguides are interconnected by flange fittings 308,309. The first waveguide 302 is protected by a sheath 310 of bio-medically inert material which is substantially transparent to microwave energy of the desired frequency. The sheath 310 is arranged to interconnect with the flange 309 so as to be removable and replaceable after each use of the

probe. The second waveguide 303 includes an excitation stub 311 which receives input from co-axial cable 312. The interconnection between the sheath 310 and the flange 309 is shown diagrammatically but will comprise a sacrifical joint causing breakage of the sheath 310 on removal, eg. it may comprise co-operating wedged ribs on the sheath 310 and the flange 309 which allow engagement but resist disengagement without breakage.

The arrangement of figures 3a, 8b and 8c employs a protective sheath 300 and a disposable handle 302 which can be supplied in a sterile pack for single use only. In order to ensure disposal of the protective sheath 300, and the handle 302 following use, the probe 301, of construction as exemplified in figure 1, is housed in the handle 302 for use. The handle 302 comprises two halves 303, 304 hinged at hinge points 305, 306. The handle 302 is moulded of microwave absorbing material and the hinged halves 303, 304 fold around the probe base and cable 307 leaving the first dielectric filled waveguide 308 and antenna portion 309 protruding from the handle as shown.

The two halves 303, 304 of the handle 302 are secured together by means of the protective sheath 300 which is placed over the protruding waveguide 308

and antenna portion 309. The sheath 300 has a sacrificial join 310 which fits over the handle halves 303, 304 and can only be removed by breaking the join 310. The sheath 300 is moulded from a biomedical material that is low-loss to microwaves.

In order to control use of the disposable handle 302 and reference the disposable items to a systems treatment log, a bar code 311 is used which can be automatically read by a bar code reader (not shown) when the assembled probe is placed in a system holster 313. The holster 313 is provided on a trolley 314 including the control elements of the system described in more detail with reference to figure 2. For example, a control keypad 315, display arm 316 and display 317 are shown.

In order to ensure that a handle 302 and sheath 300 are used with the probe 301, the cable 307 suitably includes a control switch 318 which is operative by means of a spring switch 319 on the handle 302. The control switch 313 is operative through wire 320 in the cable 307 which also includes a wire 321 from the thermocouple temperature sensor 322. The bar code 311 on the handle 302 will be unique and the software of the system is designed to reject second use to ensure disposal and replacement by a new sterile pack comprising handle and sheath

for each treatment. If desired, the sheath may also include a bar code and the bar code may include batch and date information for data logging purposes.

In most applications, and particularly in the preferred method of the invention, the probe will be used to apply heat to a load. When the load is of a biological nature, the addition of temperature sensors in the probe body as shown in some of the figures is important for safety, monitors allowing for in-situ temperature readings which can be input to feedback control and data logging systems.

In use, with reference to diagrammatic figures 9a and 9b, the probe 401 of the invention is supplied with a microwave frequency input in the region of 3-12GHz from microwave frequency generator. The dielectric material 402 within the first waveguide optimises a smooth transition without causing undue reflection. The probe 401 is suitably provided with a handle allowing manipulation by the operator and providing sterile single use as described by way of example with reference to figures 8a, 8b, 8c.

The patient is prepared by drugs being administered to contract the endometrial layer 403 of the uterus 404 as necessary. The cervix 405 is dilated and the surgeon will then insert a tool 'not shown' to determine the depth of the uterus 404 to

determine the area for treatment. The probe 401 is then inserted into the uterus 404 and the probe tip 406 positioned using markers 407 on the length of the probe as shown diagrammatically.

When the applicator tip is placed in biological tissue the generated field shape 408 in the tissue 409 can be a uniform sphere-like shape of about 4-5mm from the dielectric surface of the probe tip 406 as shown diagrammatically in figure 9a. Electromagnetic heating of the tissue 409 only occurs within this sphere.

In the particular treatment disclosed the probe 401 is inserted to the fundus of the uterus 404 and the probe 401 slowly withdrawn to expose the full endometrial lining to the electromagnetic field. The microwave electromagnetic energy radiated from the exposed probe tip 406 heats the localised area of endometrium 43 and during treatment the temperature is continually monitored by means of the temperature sensors. Thus, for example, the power may be switched on for a period of 9 seconds and then switched off for a period of 1 second whilst the temperature is measured. Whilst the control in this respect may be manual it is preferred to provide an automatic control system for maintaining the controlling temperature by means of the fibre-optic

thermometry systems and data acquisition and control means.

The microwave energy is strongly absorbed by the tissue of the endometrium and, by controlling the frequency and the power, the depth of absorption can be restricted solely to the endometrium itself which is about 5mm in depth. This has the advantage that physical injury or radiation effects on surrounding tissue are avoided. The markers 407 on the probe 401 assist the surgeon in knowing where the probe tip 406 is in the uterine cavity during treatment.

The treatment time is likely to be less than 20 minutes minimising gynaecologist time and allowing the patient a minimum time in hospital typically 1 day or less. The treated endometrium is left as scar tissue.

Although, the invention has been described using substantially continuous heating using lower power eg 50 watts to achieve a temperature in excess of 60°C, the microwave electromagnetic energy may be pulsed at a much higher power by means of a pulse magnetron. This provides pulses of kilowatt power in microseconds each pulse being spaced by the order of a millisecond. For example, it may be possible to provide pulses with a peak output of 30 kilowatts for a duration of 1 microsecond spaced by 1 millisecond.

Pulsing may have the advantage of countering the body's natural reaction to continuous heating of tissue of increasing the blood flow to the area being treated to provide cooling. Thus continuous heating may not be as efficient in destroying the cells as pulsed heating where the effect of the increased blood flow is minimised or not even promoted in the first instance.

From the drawings it will be seen that the probe of the present invention is designed to propagate and radiate microwave electromagnetic energy in a controlled fashion. The design makes use of a dielectric material within a circular waveguide with dimensions dictated by the microwave frequency used and the electrical properties of the dielectric material. The preferred dielectric material is alumina which provides an antenna diameter which is compatible with the narrow neck of the uterus. However, choosing a material with a higher dielectric constant, this diameter could be made even smaller. The dielectric material may be ceramic, plastics or other suitable material.

Although, the choice of dielectric material will fix the probe diameter, the tip of the exposed antenna portion will be shaped to achieve the desired radiation pattern. The profile of the protective

sheath can also be shaped to provide more accurate coverage of radiation in a specifically shaped load. In certain applications part of all of the probe may be designed to swivel or rotate to achieve better radiation coverage across a load. Thus, careful design of the shape and size of the probe will automatically match it to an application specific load, thus reducing the effects of standing waves which can cause loss of power and hot spots. This optimum matching can be offset by the variance of load shape and size. Tuning can be done by introducing tuning screws into the antenna/waveguide body or by adding specifically designed metal tuning washers into the dielectric/antenna assembly.

The protective sheath is, preferably of a sterile, single use, and disposable design will be used to provide a medically inert external coating for all parts of the probe that come in contact with a body. The material will be medically inert, low-loss at microwave frequencies, capable of withstanding extended exposure to harsh chemicals and high temperatures, and it will lend itself to production moulding techniques. The protective sheath suitably includes a bar code to ensure single use to prevent cross-contamination and to provide traceability.

As an alternative to bar codes, the unique identification means may comprise any other suitable means, eg. a passive electronic transponder which, if desired, may be embedded in the material of the protective sheath and/or the handle.

CLAIMS

- 1. A probe for applying electromagnetic radiation at microwave frequency to a body comprises means for receiving microwave signal input of a predetermined frequency, a waveguide for receiving and propagating said microwave frequency input, said waveguide being of a cross-sectional dimension which would not normally pass the microwaves at said frequency, dielectric material within the waveguide, the dielectric constant of which varies the cut-off frequency of the waveguide so that it may propagate desired modes of the microwaves, and an antenna portion at or adjacent an end of the probe allowing wave transmission away from the probe.
- 2. A probe according to claim 1 wherein the means for receiving the microwave signal comprises a second waveguide of larger cross-sectional dimension than the first waveguide, transition means being provided between the first and second waveguides.
- 3. A probe according to claim 2 wherein the transition means comprises a tapered waveguide interconnecting the first and second waveguide and loaded with dielectric material.
- 4. A probe according to claim 3 wherein the dielectric material in the form of a rod having a tapered end at the transition to optimise transition

and extending outwardly beyond the first waveguide to form the exposed antenna portion of the probe.

- 5. A probe according to claim 1 wherein there is a single waveguide and the means for receiving the microwave input directly excites the dielectric filled waveguide of the desired smaller cross-sectional dimension.
- 6. A probe for applying electromagnetic radiation at microwave frequency to a body comprising a first dielectric stage and a second dielectric stage, the two stages, in use, being operatively connected together,

the first dielectric stage comprising a first waveguide of a first cross-section; a second waveguide of a second cross-section larger than the cross-section of the first waveguide for receiving and propagating microwave signal input of a predetermined frequency, and transition means between the first and second waveguides including dielectric material, the dielectric constant of which varies the cut-off frequency of the first waveguide so that it may propagate said microwave signal at the predetermined frequency; and,

the second dielectric stage comprising a probe antenna of dielectric material, a third waveguide about a portion of the dielectric material and being

of substantially the same cross-section as the first waveguide, and an exposed antenna portion at or adjacent a free end of the probe allowing wave transmission away from the probe.

- 7. A probe according to claim 6 wherein the transition means of the first dielectric stage comprises a tapered waveguide interconnecting the first and second waveguides, a tapered end on the dielectric material within the tapered waveguide to optimize transition and a dielectric buffer between the tapered end of the dielectric material and the tapered waveguide, the dielectric constant of which is greater than air but less than that of the dielectric material.
- 8. A probe according to any of the preceding claims including temperature sensing means.
- 9. A probe according to any of the preceding claims including a sacrificial protective sheath adapted to be secured over the probe.
- 10. Protection means for a probe for applying electromagnetic radiation at microwave frequency to a body, said protection means comprising a sheath having a tubular body which, in use, may be passed over the probe to encapsulate the operative end of the probe and which is substantially transparent to microwaves at the intended frequency of operation.

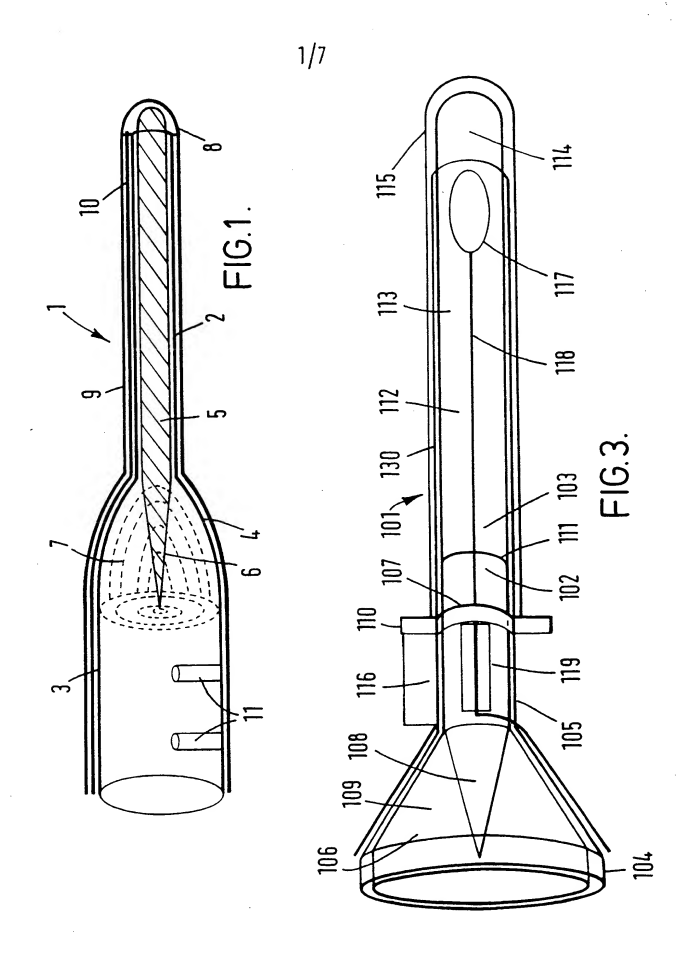
and means for securing the sheath in position whereby the sheath may be removed and discarded after use of the probe.

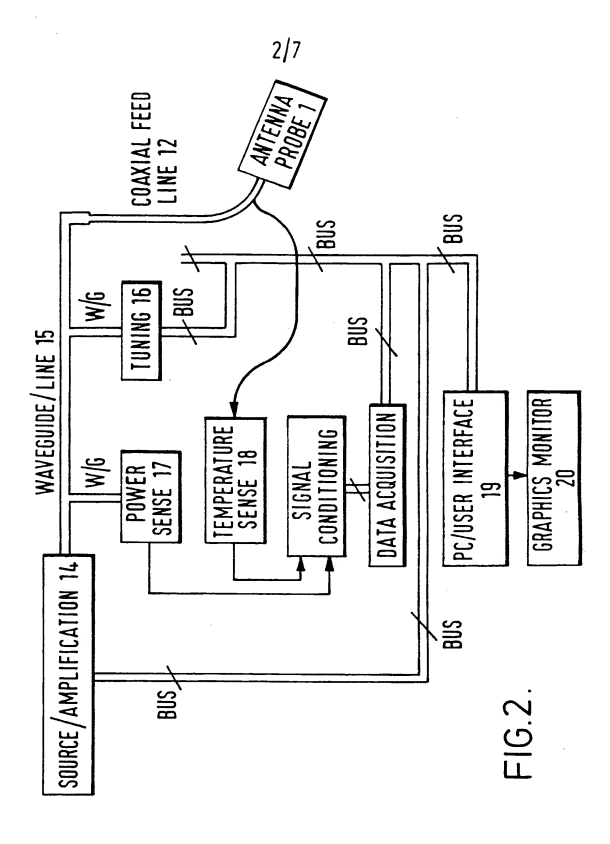
- 11. Protection means according to claim 10 further including a disposable handle arranged to receive a probe for use, the handle being locked in position about the probe by interengagement with the sheath.
- 12. Protection means according to claim 10 or 11 further including unique identification means to ensure single use.
- 13. Protection means according to claim 12 wherein the unique identification means comprises a bar code.
- 14. Protection means according to any of claims
 10 to 13 wherein the means for securing the sheath
 includes a sacrificial join ensuring single use of
 the sheath.
- 15. A sterile pack comprising a disposable handle and sheath in accordance with claim 11.
- 16. A method of endometrial ablation comprising the steps of providing a probe in accordance with any one of claims 1 to 9 having at least an operative end of outside dimensions no greater than the dimension of a dilated cervix, inserting the operative end of the probe through the cervix into the uterus.

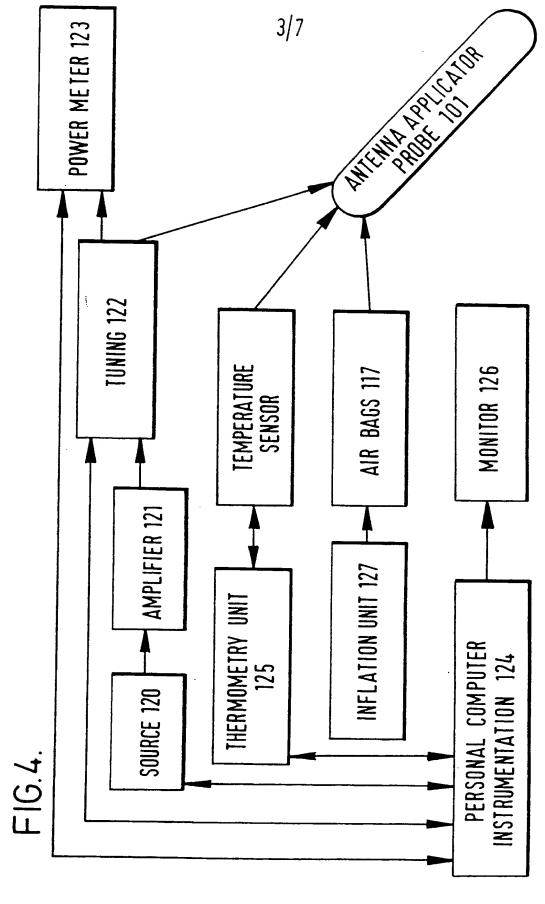
applying microwave energy to the probe at a frequency which will be substantially completely absorbed by the endometrium, monitoring the operating temperature to ensure that the endometrium tissue is heated and maintaining the application of the microwave energy for a period of time sufficient to destroy cells of the endometrium.

17. A method of endometrium ablation according to claim 16 wherein the microwave energy is applied either continually or in pulses.

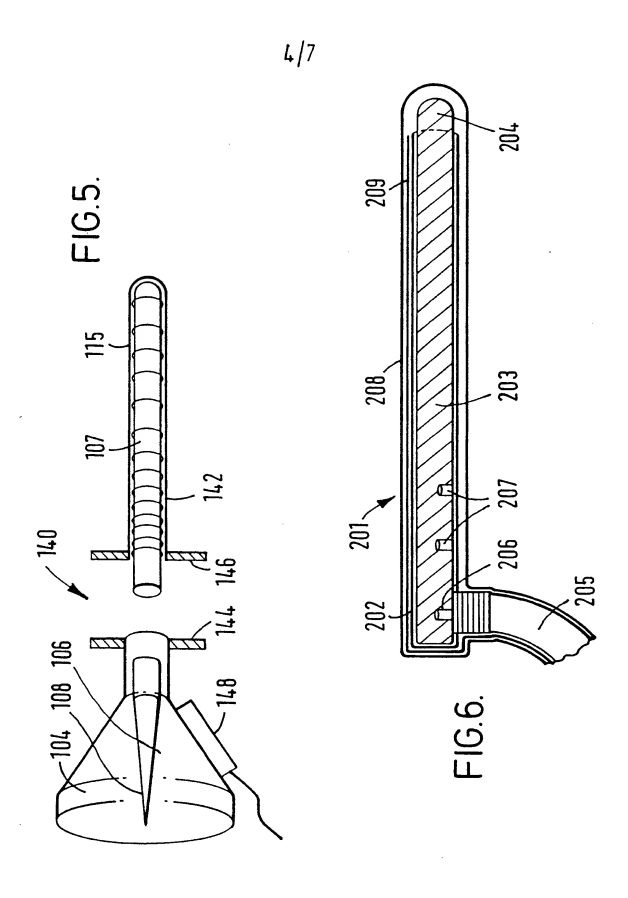
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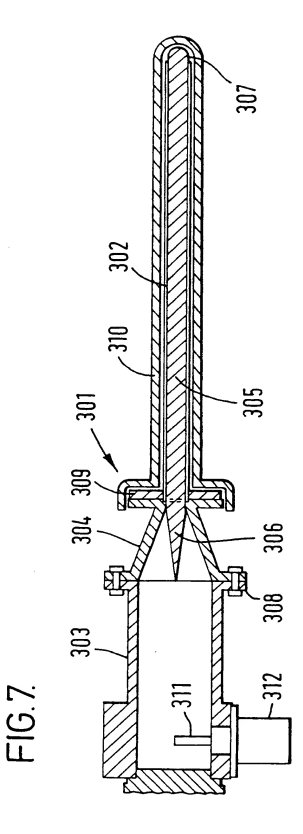


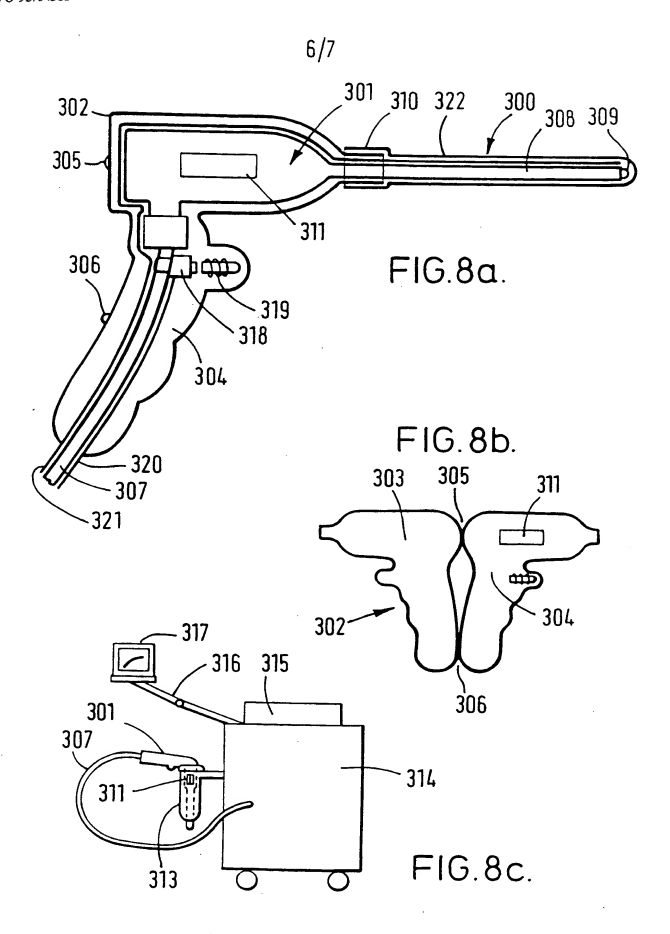


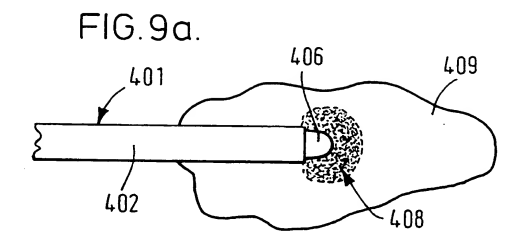
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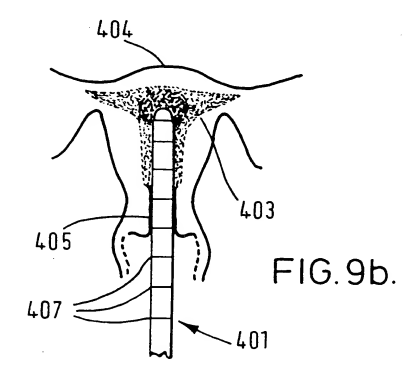


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INTE ATIONAL SEARCH REPORT

International application No. PCT/GB 94/01565

CLASSIFICATION OF SUBJECT MATTER PC 6 H01Q13/24 A61N5/ IPC 6 A61N5/04 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) H01Q A61N H01P IPC 6 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages 1-17 US,A,4 800 899 (ELLIOTT) 31 January Α see abstract; figures 1-5 1,6 PATENT ABSTRACTS OF JAPAN A vol. 6, no. 52 (E-100) (930) 7 April 1982 & JP, A, 56 166 605 (NIPPON DENSHIN) 21 December 1981 see abstract 1,6 PATENT ABSTRACTS OF JAPAN vol. 10, no. 20 (E-376) 25 January 1986 & JP,A,60 180 302 (NIPPON DENSHIN DENWA) 14 September 1985 see abstract Patent family members are listed in annex. Further documents are listed in the continuation of box C. IX I Special categories of cited documents: "I" later document published after the international filing date or priority date and not in conflict with the application but *A* document defining the general state of the art which is not considered to be of particular relevance cited to understand the principle or theory underlying the invention 'E' earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date involve an inventive step when the document is taken alone 1. document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another 'Y' document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the document is combined with one or more other such docu-ments, such combination being obvious to a person skilled document referring to an oral disclosure, use, exhibition or in the art. document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of mailing of the international search report Date of the actual completion of the international search 2 8. 11. 94 14 November 1994 Authorized officer Name and mailing address of the ISA

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